

Day : Friday Date: 8/25/2006

Time: 16:36:22

## **Inventor Name Search**

Enter the first few letters of the Inventor's Last Name. Additionally, enter the first few letters of the Inventor's First name.

Last Name	First Name	
Penttila	M	Search

To go back use Back button on your browser toolbar.



Day: Friday Date: 8/25/2006

Time: 16:36:22

### **Inventor Name Search**

Enter the first few letters of the Inventor's Last Name. Additionally, enter the first few letters of the Inventor's First name.

Last Name	First Name	
Ward	Michael	Search

To go back use Back button on your browser toolbar.



Day: Friday Date: 8/25/2006

Time: 16:36:22

## **Inventor Name Search**

Enter the first few letters of the Inventor's Last Name. Additionally, enter the first few letters of the Inventor's First name.

Last Name	First Name	
Wang	Huaming	Search

To go back use Back button on your browser toolbar.



Day: Friday Date: 8/25/2006

Time: 16:36:22

## **Inventor Name Search**

Enter the first few letters of the Inventor's Last Name. Additionally, enter the first few letters of the Inventor's First name.

Last Name	First Name	
Valkon	Ма	Search

To go back use Back button on your browser toolbar.



Day : Friday Date: 8/25/2006

Time: 16:36:22

### **Inventor Name Search**

Enter the first few letters of the Inventor's Last Name. Additionally, enter the first few letters of the Inventor's First name.

Last Name	First Name	
Salohei	Ма	Search

To go back use Back button on your browser toolbar.

# SCORE Search Results Details for Application 10663450 and Search Result us-10-663-450 5\_copy\_84\_147.rai.

Score Home Page Retrieve Application List SCORE System Overview SCORE FAQ Comments / Sugg

This page gives you Search Results detail for the Application 10663450 and Search Result us-10-665\_copy\_84\_147.rai.

start

Go Back to p

GenCore version 5.1.9
Copyright (c) 1993 - 2006 Biocceleration Ltd.

OM protein - protein search, using sw model

Run on: June 28, 2006, 01:41:45; Search time 29.44 Seconds

(without alignments)

190.284 Million cell updates/sec

Title: US-10-663-450-5\_COPY\_84\_147

Perfect score: 323

Sequence: 1 EKKPVKKRKSWGQVLPEPKT.....NRRAAQSSRERKRLEVEALE 64

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 650591 seqs, 87530628 residues

Total number of hits satisfying chosen parameters: 650591

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Issued\_Patents\_AA: \*

1: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/5\_COMB.pep:\*
2: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/6\_COMB.pep:\*
3: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/7\_COMB.pep:\*
4: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/H\_COMB.pep:\*
5: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/PCTUS\_COMB.pep:\*
6: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/PCTUS\_COMB.pep:\*

6: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/RE\_COMB.pep:\*
7: /EMC Celerra\_SIDS3/ptodata/2/iaa/backfiles1.pep:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result Query

No. Score Match Length DB ID Description

1	307	95.0	409	2	US-09-533-029-104	Sequence	104, App
2	174	53.9	248	2	US-09-248-796A-18840	Sequence	18840, A
3	99	30.7	143	2	US-09-640-211A-1930	Sequence	1930, Ap
4	89	27.6	23	2	US-09-831-642-72	Sequence	72, Appl
5	89	27.6	371	2	US-09-148-545-259	Sequence :	259, App
6	89	27.6	371	2	US-09-621-011-259	Sequence :	259, App
7	86	26.6	102	2	US-09-640-211A-1046	Sequence	1046, Ap
8	82.5	25.5	611	2	US-09-216-393B-81	Sequence	81, Appl
9	81	25.1	212	2	US-09-489-039A-13074	Sequence	13074, A
10	80.5	24.9	395	2	US-09-247-155-113	Sequence	113, App
11	80.5	24.9	395	2	US-09-513-999C-14	Sequence	14, Appl
12	80.5	24.9	395	2	US-09-471-276-14	Sequence	14, Appl
13	80.5	24.9	395	2	US-09-903-190-113	Sequence	113, App
14	80	24.8	63	2	US-09-107-433-3674	Sequence	
15	78	24.1	551	2	US-08-796-899-29	Sequence	29, Appl
16	77.5	24.0	501	2	US-09-949-016-11281	Sequence	11281, A
17	77	23.8	264	2	US-09-949-016-10789	Sequence	10789, A
18	76.5	23.7	351	2	US-09-870-089B-2	Sequence	
19	76.5	23.7	362	2	US-09-949-016-10923	Sequence	
20	75.5	23.4	463	2	US-09-345-236B-2	Sequence	
21	75	23.2	170	2	US-09-248-796A-27838	Sequence	
22	73.5	22.8	260	2	US-09-538-092-950	Sequence	
23	73.5	22.8	338	1	US-08-218-686-2	Sequence	
24	73.5	22.8	338	2	US-08-460-242-2	Sequence	<del>-</del> -
25	73.5	22.8	2058	2	US-09-949-016-6835	Sequence	
26	73.5	22.8	2111	2	US-09-949-016-10199	Sequence	_
27	73	22.6	325	2	US-09-267-031-14	Sequence	_
28	72.5	22.4	521	1	US-08-721-684C-2	Sequence	
29	72.5	22.4	521	1	US-09-005-970-2	Sequence	
30	72.5	22.4	521	2	US-09-407-715-2	Sequence	
31	71.5	22.1	307	2	US-09-267-031-6	Sequence	
32	71.5	22.1	1695	2	US-09-866-108A-15753	Sequence	
33	71	22.0	196	2	US-09-640-211A-881	Sequence	
34	71	22.0	582	2	US-09-976-594-733	Sequence	
35	71	22.0	645	2	US-09-949-016-11022	Sequence	
36	71	22.0	667	2	US-09-248-796A-14492	Sequence	
37	71	22.0	700	2	US-09-831-642-34	Sequence	
38	71	22.0	1637	2	US-09-718-692-2	Sequence	
39	71	22.0	1637	2	US-09-718-852-2	Sequence	
40	71	22.0	1637	2	US-09-718-815-2	Sequence	
41	70.5	21.8	245	2	US-09-640-211A-833	Sequence	
42	70.5	21.8	452	2	US-09-949-016-7289	Sequence	
43	70.5	21.8	472	2	US-09-520-781-22	Sequence	_
44	70.5	21.8	472	2	US-09-957-187-22	Sequence	
45	70.5	21.8	472	2	US-09-991-053-22	Sequence	
		· -		-			· -2-2-

```
RESULT 1
US-09-533-029-104
; Sequence 104, Application US/09533029
; Patent No. 6664446
; GENERAL INFORMATION:
; APPLICANT: Heard, Jacqueline
; APPLICANT: Broun, Pierre
; APPLICANT: Riechmann, Jose-Luis
; APPLICANT: Keddie, James
; APPLICANT: Pineda, Omaira
```

; APPLICANT: Adam, Luc

# **SCORE Search Results Details for Application** 10663450 and Search Result us-10-663-450-5\_copy\_84\_147.rag.

Score Home <u>Page</u>

Retrieve Application List

SCORE System Overview

**SCORE** FAO

Comments / Suggestions

This page gives you Search Results detail for the Application 10663450 and Search Result us-10-663-450-5\_copy\_84\_147.rag.

start

Go Back to previous page

GenCore version 5.1.9 Copyright (c) 1993 - 2006 Biocceleration Ltd.

OM protein - protein search, using sw model

Run on:

June 28, 2006, 01:26:15; Search time 133.12 Seconds

(without alignments)

219.816 Million cell updates/sec

Title:

US-10-663-450-5\_COPY\_84\_147

Perfect score: 323

1 EKKPVKKRKSWGQVLPEPKT.....NRRAAQSSRERKRLEVEALE 64

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched:

Sequence:

2589679 seqs, 457216429 residues

Total number of hits satisfying chosen parameters:

2589679

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

A Geneseq 8:\*

1: geneseqp1980s:\*

2: geneseqp1990s:\*

3: geneseqp2000s:\*

4: geneseqp2001s:\*
5: geneseqp2002s:\*

6: geneseqp2003as:\*

7: geneseqp2003bs:\*

8: geneseqp2004s:\*

9: geneseqp2005s:\* 10: geneseqp2006s:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

		ક્ષ				
Result		Query				
No.	Score	Match	Length	DB	ID	Description
1	323	100.0	64	5	AAE15373	Aae15373 Trichoder
2	323	100.0	450	4	AAB82975 _	Aab82975 Trichoder
3	323	100.0	451	5	AAE15371	Aae15371 Trichoder
4	323	100.0	451	8	ADS12801	Ads12801 Trichoder
5	323	100.0	451	8	ADS12804	Ads12804 Trichoder
6	307	95.0	409	7	ABO43144	Abo43144 A. thalia
7	307	95.0	409	7	ADB31925	Adb31925 Plant (A.
8	307	95.0	409	8	ADO02271	Ado02271 Thalecres
9	292	90.4	342	4	AAB82977	Aab82977 Aspergill
10	292	90.4	342	5	AAE15381	Aae15381 Aspergill
11	292	90.4	342	8	ADS12818	Ads12818 Aspergill
12	292	90.4	386	5	AAE15379	Aae15379 Aspergill
13	292	90.4	386	8	ADS12815	Ads12815 Aspergill
14	288	89.2	64	5	AAE15374	Aae15374 Aspergill
15	288	89.2	349	8	ADS12805	Ads12805 Aspergill
16	288	89.2	349	8	ADS12803	Ads12803 Aspergill
17	288	89.2	349	8	ADS12863	Ads12863 Aspergill
18	288	89.2	350	4	AAB82976	Aab82976 Aspergill
19	288	89.2	350	5	AAE15372	Aae15372 Aspergill
20	175	54.2	68	5	AAE15382	Aae15382 Yeast HAC
21	175	54.2	68	8	ADS12859	Ads12859 Saccharom
22	175	54.2	84	5	ABP02534	Abp02534 Human ORF
23	175	54.2	200	8	ADS43437	Ads43437 Bacterial
24	175	54.2	230	2	AAW53806	Aaw53806 Transcrip
25	175	54.2	230	8	ADT87049	Adt87049 Yeast Str
26	175	54.2	238	2	AAW53807	Aaw53807 Transcrip
27	108	33.4	192	4	AAB82614	Aab82614 Maize roo
28	108	33.4	192	4	AAB82615	Aab82615 Maize roo
29	108	33.4	192	4	AAB82616	Aab82616 Maize roo
3 0	108	33.4	192	4	AAG66525	Aag66525 Maize roo
31	108	33.4	192	4	AAG66526	Aag66526 Maize roo
32	106	32.8	168	5	AAU93013	Aau93013 Arabidops
33	106	32.8	168	7	ADD30174	Add30174 Plant yie
34	106	32.8	168	8	ADI43893	Adi43893 Plant tra
35	106	32.8	211	3	AAG08861	Aag08861 Arabidops
36	103	31.9	170	8	ADM48147	Adm48147 Polypepti
37	101	31.3	672	8	ADX95805	Adx95805 Plant ful
38	100.5	31.1	185	9	ADW17162	Adw17162 Eucalyptu
39	99	30.7	143	3	AAB33151	Aab33151 Pinus rad
40	99	30.7	163	9	ADW17580	Adw17580 Pinus rad
41	94	29.1	646	7	ABM86095	Abm86095 Rice abio
42	93	28.8	120	3	AAG27808	Aag27808 Arabidops
43	93	28.8	135	3	AAG07181	Aag07181 Arabidops
44	93	28.8	149	3	AAG07181 AAG07180	Aag07180 Arabidops
45	93	28.8	149	3	AAG27807	Aag27807 Arabidops
4.7	23	20.0	747	ی	123G2 / GU /	nag2/00/ Arabidops

```
RESULT 1
    AAE15373 standard; protein; 64 AA.
XX
AC
    AAE15373;
XX
DT
    29-AUG-2003 (revised)
```

# SCORE Search Results Details for Application 10663450 and Search Result us-10-663-450-19\_copy\_45\_116.rup.

Score Home Page Retrieve Application List SCORE System Overview

SCORE FAO Comments / Suggestions

This page gives you Search Results detail for the Application 10663450 and Search Result us-10-663-450-19\_copy\_45\_116.rup.

<u>start</u>

Go Back to previous page

GenCore version 5.1.9
Copyright (c) 1993 - 2006 Biocceleration Ltd.

OM protein - protein search, using sw model

Run on:

June 28, 2006, 01:26:19; Search time 169.56 Seconds

(without alignments)

392.788 Million cell updates/sec

Title:

US-10-663-450-19 COPY 45 116

Perfect score: 365

Sequence: 1 EEKKPVKKRKSWGQELPVPK.....SRERKRLEMEKLENEKIQME 72

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2849598 seqs, 925015592 residues

Total number of hits satisfying chosen parameters: 2849598

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

UniProt\_7.2:\*

1: uniprot\_sprot:\*
2: uniprot\_trembl:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

#### SUMMARIES

Result No.	Score	% Query Match	Length	DB	ID	Description
1	365	100.0	342	2	Q6W8X2_ASPNG	Q6w8x2 aspergillus
2	361	98.9	348	2	Q2U743_ASPOR	Q2u743 aspergillus
3	358	98.1	433	2	Q4WEY8_ASPFU	Q4wey8 aspergillus
4	351	96.2	347	2	Q5AQN3 EMENI	Q5aqn3 aspergillus

5	351	96.2	350	2	Q8TFU8_EMENI		emericella
6	298	81.6	451	2	Q8TFF3_TRIRE	Q8tff3	trichoderma
7	297	81.4	430	2	Q7SHF0_NEUCR	<del>-</del>	neurospora
8	270	74.0	556	2	Q2KH12_MAGGR	Q2kh12	magnaporthe
9	261	71.5	429	2	Q4HTT5_GIBZE	Q4htt5	gibberella
10	179.5	49.2	299	2	Q6CEV1_YARLI	Q6cev1	yarrowia li
11	177	48.5	357	2	Q5AA52_CANAL	Q5aa52	candida alb
12	175.5	48.1	273	2	Q6CKQ1_KLULA	Q6ckq1	kluyveromyc
13	174	47.7	260	2	Q6BQC2_DEBHA	Q6bqc2	debaryomyce
14	172.5	47.3	230	1	HAC1_YEAST	P41546	saccharomyc
15	168	46.0	228	2	Q75BQ5_ASHGO	Q75bq5	ashbya goss
16	145	39.7	329	2	Q6FLY3 CANGA	Q6fly3	candida ġla
17	123	33.7	168	1	HY5_ARATH	024646	arabidopsis
18	118	32.3	336	2	Q5VR11_ORYSA	Q5vrll	oryza sativ
19	116	31.8	69	2	Q58L60_BRACM	Q58160	brassica ca
20	116	31.8	109	2	Q84XX8_BRARP	Q84xx8	brassica ra
21	114.5	31.4	252	2	Q6NX18_XENTR	Q6nx18	xenopus tro
22	113.5	31.1	254	2	Q2TAU5_XENLA	Q2tau5	xenopus lae
23	113.5	31.1	350	2	Q90ZR7_XENLA	Q90zr7	xenopus lae
24	113.5	31.1	396	2	Q7ZYC2_XENLA	Q7zyc2	xenopus lae
25	107	29.3	176	2	Q69XK6_ORYSA	Q69xk6	oryza sativ
26	106	29.0	261	2	Q3SZZ2_BOVIN	Q3szz2	bos taurus
27	105	28.8	158	1	HY5 LYCES	Q9sm50	lycopersico
28	105	28.8	703	2	Q4WC74_ASPFU	Q4wc74	aspergillus
29	104	28.5	627	2	Q5BD44 EMENI	Q5bd44	aspergillus
30	103.5	28.4	208	2	Q5DFK2_SCHJA	Q5dfk2	schistosoma
31	103	28.2	263	2	Q8UVQ5_BRARE	Q8uvq5	brachydanio
32	103	28.2	263	2	Q90X27_BRARE	Q90x27	brachydanio
33	103	28.2	383	2	Q8QHJ5_BRARE	Q8qhj5	brachydanio
34	103	28.2	383	2	Q90XD3_BRARE	Q90xd3	brachydanio
35	102.5	28.1	141	2	Q677A7_HYAOR	Q677a7	hyacinthus
36	102.5	28.1	260	2	Q6EZA7 OREMO	Q6eza7	oreochromis
37	102.5	28.1	260	2	Q6EZA8_OREMO	Q6eza8	oreochromis
38	102.5	28.1	261	2	Q3LFR3_FUGRU	Q3lfr3	fugu rubrip
39	102	27.9	321	2	Q8GRY7_LOTJA	Q8gry7	lotus japon
40	101.5	27.8	309	2	Q4H2M2_CIOIN	Q4h2m2	ciona intes
41	101.5	27.8	385	2	Q4RTW2_TETNG	Q4rtw2	tetraodon n
42	101	27.7	261	1	XBP1_HUMAN		homo sapien
43	100	27.4	267	1	XBP1_RAT	Q9r1s4	rattus norv
44	99	27.1	188	2	Q6ZHT8_ORYSA	Q6zht8	oryza sativ
45	98	26.8	267	1	XBP1_MOUSE	035426	mus musculu

```
RESULT 1
Q6W8X2 ASPNG
    Q6W8X2_ASPNG PRELIMINARY;
                                   PRT;
                                          342 AA.
AC
    Q6W8X2;
    05-JUL-2004, integrated into UniProtKB/TrEMBL.
    05-JUL-2004, sequence version 1.
DT
    07-FEB-2006, entry version 9.
DT
DΕ
    Transcription factor HACA.
GN
    Name=hacA;
os
    Aspergillus niger.
OC
     Eukaryota; Fungi; Ascomycota; Pezizomycotina; Eurotiomycetes;
     Eurotiales; Trichocomaceae; mitosporic Trichocomaceae; Aspergillus.
OC
ОХ
     NCBI_TaxID=5061;
RN
     [1]
RΡ
     NUCLEOTIDE SEQUENCE.
```

### Score Home Page Retrieve Application List SCORE System Overview SCORE FAQ Comments / Sugg

This page gives you Search Results detail for the Application 10663450 and Search Result us-10-66 start

> GenCore version 5.1.9 Copyright (c) 1993 - 2006 Biocceleration Ltd.

OM protein - protein search, using sw model

June 28, 2006, 01:33:51; Search time 20.88 Seconds

(without alignments)

331.782 Million cell updates/sec

Title: US-10-663-450-19 COPY 45 116

Perfect score: 365

Sequence: 1 EEKKPVKKRKSWGQELPVPK.....SRERKRLEMEKLENEKIQME 72

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 283416 seqs, 96216763 residues

Total number of hits satisfying chosen parameters: 283416

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : PIR 80:\*

> 1: pir1:\* 2: pir2:\*

3: pir3:\*

4: pir4:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

#### SUMMARIES

			0				
R	esult		Query				
	No.	Score	Match	Length	DB	ID	Description
	1	172.5	47.3	238	2	s78571	transcription fact
	2	123	33.7	168	2	T50922	bZIP protein HY5 [
	3	100	27.4	267	· 2	JC4857	hepatocarcinogenes
	4	98	26.8	266	. 2	JC7300	tax-responsive ele
	5	98	26.8	322	2	T08592	TGACG-motif-bindin
	6	98	26.8	326	2	T08591	TGACG-motif bindin
	7	97.5	26.7	260	1	A36299	transcription fact
	8	95	26.0	322	2	T12093	TGACG-motif bindin

9	94.5	25.9	506	2	D84609	hypothetical prote
10	89.5	24.5	600	2	T00759	hypothetical prote
11	88.5	24.2	452	2	Н96710	hypothetical prote
12	88	24.1	338	1	TVMSFB	transforming prote
13	88	24.1	338	2	153043	transforming prote
14	87.5	24.0	176	2	B90087	hypothetical prote
15	87.5	24.0	737	2	T15597	hypothetical prote
16	87.5	24.0	849	1	S64732	scaffold attachmen
17	87.5	24.0	1017	2	T15598	hypothetical prote
18	87	23.8	1549	1	A40691	trichohyalin - she
19	85.5	23.4	688	2	T32750	hypothetical prote
20	85	23.3	468	2	S33222	transcription fact
21	85	23.3	1240	2	S52734	hypothetical prote
22	84.5	23.2	246	2	T12585	Dc3 promoter-bindi
23	83.5	22.9	486	2	JC4028	activating transcr
24	82.5	22.6	332	2	S15347	transcription fact
25	82.5	22.6	351	2	A45377	transcription fact
26	82.5	22.6	360	2	т03373	probable G-box bin
27	82.5	22.6	515	2	A42140	box B-binding fact
28	82.5	22.6	516	2	A44494	cAMP-responsive el
29	82	22.5	381	2	S26812	transcription fact
30	81.5	22.3	774	2	S25284	protein kinase nek
31	81	22.2	349	2	A41349	histone-specific t
32	81	22.2	349	2	S77570	transcription fact
33	81	22.2	433	2	JC1230	DNA-binding protei
34	80.5	22.1	839	2	150590	class I INCENP pro
35	80.5	22.1	877	2	<b>I50591</b>	class II INCENP pr
36	80	21.9	313	2	A34785	DNA-binding protei
37	80	21.9	331	2	S33223	transcription fact
38	80	21.9	358	2	C42026	cyclic AMP respons
39	80	21.9	389	1	A39429	cAMP response elem
40	80	21.9	448	2	A42026	cAMP response elem
41	80	21.9	456	2	B42026	cyclic AMP respons
42	80	21.9	505	1	S05380	transcription fact
43	80	21.9	771	1	A33430	h-caldesmon - chic
44	79.5	21.8	372	2	S17715	transcription acti
45	79.5	21.8	483	2	S12741	transcription fact

```
ALIGNMENTS
RESULT 1
S78571
transcription factor HAC1 - yeast (Saccharomyces cerevisiae)
N; Alternate names: protein YFL031w
C; Species: Saccharomyces cerevisiae
C;Date: 13-Sep-1998 #sequence_revision 13-Sep-1998 #text change 09-Jul-2004
C; Accession: S78571; S56223; S53578
R; Murakami, Y.
submitted to the Protein Sequence Database, January 1998
A; Reference number: $78570
A; Accession: S78571
A; Molecule type: DNA
A; Residues: 1-238
A; Cross-references: UNIPROT: P41546; UNIPARC: UPI0000157F61; EMBL: D50617; MIPS: YFL031w
A; Note: this is a revision to the sequence from reference S56186
R; Murakami, Y.; Naitou, M.; Hagiwara, H.; Shibata, T.; Ozawa, M.; Sasanuma, S.I.; Sasa
submitted to the EMBL Data Library, May 1995
A; Description: Analysis of the nucleotide sequence of chromosome VI from Saccaromyces
```

A; Reference number: S56186

# SCORE Search Results Details for Application 10663450 and Search Result us-10-663-450-19\_copy\_45\_116.rapbn.

Score Home Page Retrieve Application List

SCORE System Overview

SCORE FAQ

Comments / Suggestions

This page gives you Search Results detail for the Application 10663450 and Search Result us-10-663-450-19\_copy\_45\_116.rapbn.

<u>start</u>

Go Back to previous pag

GenCore version 5.1.9
Copyright (c) 1993 - 2006 Biocceleration Ltd.

OM protein - protein search, using sw model

Run on:

June 28, 2006, 01:44:20 ; Search time 9.72 Seconds

(without alignments)

169.226 Million cell updates/sec

Title:

US-10-663-450-19\_COPY 45 116

Perfect score: 365

Sequence:

1 EEKKPVKKRKSWGQELPVPK.....SRERKRLEMEKLENEKIQME 72

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched:

99297 seqs, 22845552 residues

Total number of hits satisfying chosen parameters: 99297

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

Published\_Applications\_AA New:\*

1: /EMC\_Celerra\_SIDS3/ptodata/1/pubpaa/US09\_NEW\_PUB.pep:\*

2: /EMC\_Celerra\_SIDS3/ptodata/1/pubpaa/US06\_NEW\_PUB.pep:\*

3: /EMC\_Celerra\_SIDS3/ptodata/1/pubpaa/US07\_NEW\_PUB.pep:\*

4: /EMC\_Celerra\_SIDS3/ptodata/1/pubpaa/US08 NEW PUB.pep:\*

5: /EMC\_Celerra\_SIDS3/ptodata/1/pubpaa/vS08\_NEW\_PUB.pep:\*

6: /EMC\_Celerra\_SIDS3/ptodata/1/pubpaa/US10\_NEW\_PUB.pep:\*

7: /EMC\_Celerra\_SIDS3/ptodata/1/pubpaa/US11 NEW PUB.pep:\*

8: /EMC\_Celerra\_SIDS3/ptodata/1/pubpaa/US60\_NEW\_PUB.pep:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result

Query

No.	Score	Match	Length	DB	ID	Description
1	275	75.3	425	6	US-10-449-902-40262	Sequence 40262, A
2	108	29.6	134	6	US-10-953-349-13422	Sequence 13422, A
3	108	29.6	134	6	US-10-953-349-22329	Sequence 22329, A
4	108	29.6	191	6	US-10-953-349-13421	Sequence 13421, A
5	108	29.6	191	6	US-10-953-349-22328	Sequence 22328, A
6	98	26.8	199	6	US-10-953-349-12209	Sequence 12209, A
7	98	26.8	201	6	US-10-953-349-20562	Sequence 20562, A
8	98	26.8	318	6	US-10-953-349-12208	Sequence 12208, A
9	98	26.8	320	6	US-10-953-349-20561	Sequence 20561, A
10	98	26.8	324	6	US-10-953-349-12207	Sequence 12207, A
11	98	26.8	326	6	US-10-953-349-20560	Sequence 20560, A
12	95.5	26.2	646	6	US-10-449-902-51369	Sequence 51369, A
13 14	90.5 90.5	24.8 24.8	293 318	6 6	US-10-953-349-23767	Sequence 23767, A
15	90.5	24.8	323	6	US-10-953-349-23766	Sequence 23766, A
16	89.5	24.5	335	6	US-10-953-349-23765 US-10-449-902-34705	Sequence 23765, A Sequence 34705, A
17	88	24.1	467	7	US-11-293-697-3606	-
18	88	24.1	611	7	US-11-321-421-81	Sequence 3606, Ap Sequence 81, Appl
19	87.5	24.0	675	7	US-11-293-697-3326	Sequence 3326, Ap
20	81.5	22.3	296	6	US-10-953-349-24027	Sequence 24027, A
21	81.5	22.3	362	6	US-10-953-349-24026	Sequence 24026, A
22	81.5	22.3	382	6	US-10-953-349-24025	Sequence 24025, A
23	81.5	22.3	467	6	US-10-449-902-38097	Sequence 38097, A
24	81	22.2	332	6	US-10-953-349-23520	Sequence 23520, A
25	81	22.2	457	6	US-10-449-902-53115	Sequence 53115, A
26	81	22.2	523	6	US-10-449-902-56056	Sequence 56056, A
27	81	22.2	650	6	US-10-449-902-44708	Sequence 44708, A
28	80	21.9	142	6	US-10-449-902-31867	Sequence 31867, A
29	79.5	21.8	675	6	US-10-449-902-42244	Sequence 42244, A
30	78.5	21.5	539	6	US-10-449-902-54234	Sequence 54234, A
31	78	21.4	147	6	US-10-953-349-36806	Sequence 36806, A
32 33	78 78	21.4	165 186	6 6	US-10-953-349-36805	Sequence 36805, A
34	77.5	21.4	437	6	US-10-953-349-36804 US-10-449-902-40619	Sequence 36804, A
35	77.3	21.1	. 484	6	US-10-449-902-47431	Sequence 40619, A Sequence 47431, A
36	77	21.1	484	6	US-10-449-902-55580	Sequence 55580, A
37	76	20.8	215	6	US-10-449-902-45271	Sequence 45271, A
38	76	20.8	301	6	US-10-449-902-43842	Sequence 43842, A
39	75.5	20.7	281	6	US-10-971-483-2	Sequence 2, Appli
40	75.5	20.7	357	6	US-10-449-902-44625	Sequence 44625, A
41	75.5	20.7	380	6	US-10-449-902-43338	Sequence 43338, A
42	75.5	20.7	577	7	US-11-289-102-220	Sequence 220, App
43	75	20.5	167	6	US-10-449-902-40094	Sequence 40094, A
44	75	20.5	253	6	US-10-953-349-11914	Sequence 11914, A
45	75	20.5	334	6	US-10-449-902-53341	Sequence 53341, A

#### RESULT 1

US-10-449-902-40262

- ; Sequence 40262, Application US/10449902
- ; Publication No. US20060123505A1
- ; GENERAL INFORMATION:
- ; APPLICANT: National Institute of Agrobiological Sciences.
- ; APPLICANT: Bio-oriented Technology Research Advancement Institution.
- ; APPLICANT: The Institute of Physical and Chemical Research.
- ; APPLICANT: Foundation for Advancement of International Science.

# SCORE Search Results Details for Application 10663450 and Search Result us-10-663-450-19\_copy\_45\_116.rapbm.

Score Home <u>Page</u>

Retrieve Application SCORE System

List

Overview

**SCORE** FAQ

Comments / Suggestions

This page gives you Search Results detail for the Application 10663450 and Search Result us-10-663-450-19\_copy\_45\_116.rapbm.

start

Go Back to previous page

GenCore version 5.1.9 Copyright (c) 1993 - 2006 Biocceleration Ltd.

OM protein - protein search, using sw model

Run on:

June 28, 2006, 01:42:39; Search time 113.4 Seconds

(without alignments)

294.105 Million cell updates/sec

US-10-663-450-19 COPY 45 116

Perfect score: 365

Sequence:

1 EEKKPVKKRKSWGQELPVPK.....SRERKRLEMEKLENEKIOME 72

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched:

2097797 segs, 463214858 residues

Total number of hits satisfying chosen parameters:

2097797

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

Published Applications AA Main:\*

1: /EMC\_Celerra SIDS3/ptodata/2/pubpaa/US07\_PUBCOMB.pep:\*

2: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US08\_PUBCOMB.pep:\*

3: /EMC\_Celerra SIDS3/ptodata/2/pubpaa/US09 PUBCOMB.pep:\*

4: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US10A\_PUBCOMB.pep:\*

5: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US10B\_PUBCOMB.pep:\*

6: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US11\_PUBCOMB.pep:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

#### SUMMARIES

Result

Query

No. Score Match Length DB ID

```
1
       365 100.0
                    342 3 US-09-816-277-19
                                                       Sequence 19, Appl
 2
       365
           100.0
                    342 4
                            US-10-663-450-19
                                                       Sequence 19, Appl
 3
       365 100.0
                    386 3 US-09-816-277-16
                                                       Sequence 16, Appl
 4
       365 100.0
                    386 4 US-10-663-450-16
                                                       Sequence 16, Appl
 5
       351
            96.2
                    349 3 US-09-816-277-4
                                                       Sequence 4, Appli
 6
       351
            96.2
                    349 3 US-09-816-277-6
                                                       Sequence 6, Appli
 7
                    349 4 US-10-663-450-4
       351
            96.2
                                                       Sequence 4, Appli
 8
       351
                    349 4 US-10-663-450-6
             96.2
                                                       Sequence 6, Appli
                    451 3 US-09-816-277-2
 9
       298
            81.6
                                                       Sequence 2, Appli
10
       298
            81.6
                    451 3 US-09-816-277-5
                                                       Sequence 5, Appli
       298
11
            81.6
                    451 4 US-10-663-450-2
                                                       Sequence 2, Appli
12
       298
            81.6
                    451 4 US-10-663-450-5
                                                       Sequence 5, Appli
      295
13
            80.8
                    409 3 US-09-533-029-104
                                                       Sequence 104, App
14
      295
            80.8
                    409 4 US-10-295-403-158
                                                       Sequence 158, App
15
      295
            80.8
                    409 4 US-10-412-699B-684
                                                       Sequence 684, App
16
       271
            74.2
                    174 4 US-10-767-701-61238
                                                       Sequence 61238, A
17
    172.5
            47.3
                    200 4 US-10-369-493-21867
                                                       Sequence 21867, A
18
      171
            46.8
                    68 3 US-09-816-277-60
                                                       Sequence 60, Appl
                    68 4 US-10-663-450-60
19
      171
            46.8
                                                       Sequence 60, Appl
                    168 3 US-09-934-455-102
20
      123
            33.7
                                                       Sequence 102, App
21
                    168 4 US-10-225-066A-206
      123
            33.7
                                                       Sequence 206, App
                    168 4 US-10-374-780A-2356
22
      123
            33.7
                                                       Sequence 2356, Ap
23
      123
            33.7
                    168 5 US-10-732-923-13692
                                                       Sequence 13692, A
24
      123
            33.7
                    168 5 US-10-225-066A-206
                                                       Sequence 206, App
25
      118
            32.3
                    203 4 US-10-437-963-151695
                                                       Sequence 151695,
26
      118
            32.3
                    203 5 US-10-732-923-13694
                                                       Sequence 13694, A
27
      118
            32.3
                    203 5 US-10-732-923-13695
                                                       Sequence 13695, A
28
      116
            31.8
                    109 5 US-10-732-923-13697
                                                       Sequence 13697, A
29
      113
            31.0
                    147 4 US-10-425-115-187302
                                                       Sequence 187302,
30
    110.5
            30.3
                    192 3 US-09-772-656-2
                                                       Sequence 2, Appli
                    192 3 US-09-772-656-6
31
    110.5
            30.3
                                                       Sequence 6, Appli
    110.5
                    192 3 US-09-772-656-10
32
            30.3
                                                       Sequence 10, Appl
33
    110.5
                    192 4 US-10-396-199A-2
            30.3
                                                      Sequence 2, Appli
                    192 4 US-10-396-199A-6
    110.5
34
            30.3
                                                      Sequence 6, Appli
                    192 4 US-10-396-199A-10
    110.5
35
            30.3
                                                      Sequence 10, Appl
                    143 5 US-10-856-499-1930
            29.9
36
      109
                                                      Sequence 1930, Ap
                    196 4 US-10-425-115-197786
37
      108
            29.6
                                                      Sequence 197786,
                    176 4 US-10-437-963-164243
38
      107
            29.3
                                                      Sequence 164243,
39
    105.5
            28.9
                    170 4 US-10-310-154-565
                                                      Sequence 565, App
            28.6
40
    104.5
                    168 4 US-10-425-115-246428
                                                      Sequence 246428,
41
    103.5
            28.4
                    191 4 US-10-424-599-177263
                                                      Sequence 177263,
42.
            28.2
      103
                    181 5 US-10-732-923-450
                                                      Sequence 450, App
43
    101.5
            27.8
                    114 4 US-10-425-115-246430
                                                      Sequence 246430,
44
      101
            27.7
                    217 4 US-10-425-115-197783
                                                      Sequence 197783,
45
      101
            27.7
                    261 4 US-10-205-823-453
                                                      Sequence 453, App
```

```
RESULT 1
US-09-816-277-19
; Sequence 19, Application US/09816277
; Publication No. US20010034045A1
; GENERAL INFORMATION:
; APPLICANT: Penttila, Merja E.
; APPLICANT: Ward, Michael
; APPLICANT: Wang, Huaming
; APPLICANT: Valkonen, Mari J.
; APPLICANT: Saloheimo, Markku
; TITLE OF INVENTION: Increased Production of Secreted
```

# **SCORE Search Results Details for Application** 10663450 and Search Result us-10-663-450-19\_copy\_45\_116.rai.

Score Home Page

**Retrieve Application** 

List

SCORE System Overview

SCORE FAQ

Comments / Suggestions

This page gives you Search Results detail for the Application 10663450 and Search Result us-10-66 450-19\_copy\_45\_116.rai.

start

Go Back to previous

GenCore version 5.1.9 Copyright (c) 1993 - 2006 Biocceleration Ltd.

OM protein - protein search, using sw model

Run on:

June 28, 2006, 01:41:45; Search time 33.12 Seconds

(without alignments)

190.284 Million cell updates/sec

US-10-663-450-19 COPY 45 116

Perfect score: 365

Sequence:

1 EEKKPVKKRKSWGQELPVPK......SRERKRLEMEKLENEKIQME 72

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched:

650591 seqs, 87530628 residues

Total number of hits satisfying chosen parameters:

650591

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

Issued Patents AA:\*

1: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/5\_COMB.pep:\*

2: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/6\_COMB.pep:\*

3: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/7\_COMB.pep:\*

4: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/H\_COMB.pep:\*

5: /EMC\_Celerra SIDS3/ptodata/2/iaa/PCTUS COMB.pep:\*

6: /EMC Celerra SIDS3/ptodata/2/iaa/RE\_COMB.pep:\*

7: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/backfiles1.pep:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result

Query

Score Match Length DB

Description

1	295	80.8	409	2	US-09-533-029-104	Sequence 104, App
2	177	48.5	248	2	US-09-248-796A-18840	Sequence 18840, A
3	109	29.9	143	2	US-09-640-211A-1930	Sequence 1930, Ap
4	101	27.7	264	2	US-09-949-016-10789	Sequence 10789, A
5	100.5	27.5	102	2	US-09-640-211A-1046	Sequence 1046, Ap
6	97.5	26.7	260	2	US-09-538-092-950	Sequence 950, App
7	88	24.1	338	1	US-08-218-686-2	Sequence 2, Appli
8	88	24.1	338	2	US-08-460-242-2	Sequence 2, Appli
9	88	24.1	452	2	US-09-949-016-7289	Sequence 7289, Ap
10	88	24.1	611	2	US-09-216-393B-81	Sequence 81, Appl
11	88	24.1	1637	2	US-09-718-692-2	Sequence 2, Appli
12	88	24.1	1637	2	US-09-718-852-2	Sequence 2, Appli
13	88	24.1	1637	2	US-09-718-815-2	Sequence 2, Appli
14	87.5	24.0	931	2	US-09-949-016-9850	Sequence 9850, Ap
. 15	85.5	23.4	472	2	US-09-520-781-22	Sequence 22, Appl
16	85.5	23.4	472	2	US-09-957-187-22	Sequence 22, Appl
17	85.5	23.4	472	2	US-09-991-053-22	Sequence 22, Appl
18	85.5	23.4	491	2	US-09-520-781-24	Sequence 24, Appl
19	85.5	23.4	491	2	US-09-957-187-24	Sequence 24, Appl
20	85.5	23.4	491	2	US-09-991-053-24	Sequence 24, Appl
21	85.5	23.4	521	1	US-08-721-684C-2	Sequence 2, Appli
22	85.5	23.4	521	1	US-09-005-970-2	Sequence 2, Appli
23	85.5	23.4	521	2	US-09-407-715-2	Sequence 2, Appli
24	85	23.3	1240	2	US-09-538-092-658	Sequence 658, App
25	84	23.0	667	2	US-09-248-796A-14492	Sequence 14492, A
26	83	22.7	23	2	US-09-831-642-72	Sequence 72, Appl
27	82.5	22.6	307	2	US-09-267-031-6	Sequence 6, Appli
28	82.5	22.6	351	2	US-09-870-089B-2	Sequence 2, Appli
29	82.5	22.6	362	2	US-09-949-016-10923	Sequence 10923, A
30	81.5	22.3	118	2	US-09-134-001C-2856	Sequence 2856, Ap
31	81.5	22.3	616	2	US-09-873-404-4	Sequence 4, Appli
32	81.5	22.3	616	2	US-10-243-735-4	Sequence 4, Appli
33	81.5	22.3	616	2	US-10-730-010-4	Sequence 4, Appli
34	81.5	22.3	802	2	US-09-823-240A-2	Sequence 2, Appli
35	81	22.2	422	2	US-09-248-796A-19321	Sequence 19321, A
36	80.5	22.1	470	2	US-10-104-047-3775	Sequence 3775, Ap
37	80.5	22.1	483	2	US-09-949-016-7773	Sequence 7773, Ap
38	80.5	22.1	603	2	US-10-094-749-1914	Sequence 1914, Ap
39	80.5	22.1	1034	2	US-09-976-594-590	Sequence 590, App
40	80	21.9	63	2	US-09-107-433-3674	Sequence 3674, Ap
41	80	21.9	501	2	US-09-949-016-11281	Sequence 11281, A
42	79.5	21.8	235	2	US-09-270-767-36681	Sequence 36681, A
43	79.5	21.8	235	2	US-09-270-767-51898	Sequence 51898, A
44	79 70	21.6	170	2	US-09-248-796A-27838	Sequence 27838, A
45	79	21.6	325	2	US-09-267-031-14	Sequence 14, Appl

```
RESULT 1
US-09-533-029-104
; Sequence 104, Application US/09533029
; Patent No. 6664446
; GENERAL INFORMATION:
; APPLICANT: Heard, Jacqueline
; APPLICANT: Broun, Pierre
; APPLICANT: Riechmann, Jose-Luis
```

; APPLICANT: Keddie, James ; APPLICANT: Pineda, Omaira

# **SCORE Search Results Details for Application** 10663450 and Search Result us-10-663-450copy\_45\_116.rag.

Score Home Page

Retrieve Application

**SCORE System Overview** 

**SCORE** FAQ

Suggestions

This page gives you Search Results detail for the Application 10663450 and Search Result us-10-663-450-19\_copy\_45\_116.rag.

start

) a

Go Back to previous page

GenCore version 5.1.9 Copyright (c) 1993 - 2006 Biocceleration Ltd.

OM protein - protein search, using sw model

Run on:

June 28, 2006, 01:26:15; Search time 149.76 Seconds

(without alignments)

219.816 Million cell updates/sec

US-10-663-450-19 COPY 45 116

Perfect score: 365

Sequence:

1 EEKKPVKKRKSWGQELPVPK.....SRERKRLEMEKLENEKIQME 72

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched:

2589679 seqs, 457216429 residues

Total number of hits satisfying chosen parameters: 2589679

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

A\_Geneseq\_8:\*

1: geneseqp1980s:\*

geneseqp1990s:\*

3: geneseqp2000s:\* 4: geneseqp2001s:\*

5: geneseqp2002s:\*

6: geneseqp2003as:\*

7: geneseqp2003bs:\*

8: geneseqp2004s:\*

9: geneseqp2005s:\*

10: geneseqp2006s:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result		* ^******				
No.	Caara	Query	Tanabh	חח	TD	
NO.	Score	Macch	Length		ID	Description
1	365	100.0	342	4	AAB82977	Aab82977 Aspergill
2	365	100.0	342	5	AAE15381	Aae15381 Aspergill
3	365	100.0	342	8	ADS12818	Ads12818 Aspergill
4	365	100.0	386	5	AAE15379	Aae15379 Aspergill
5	365	100.0	386	8	ADS12815	Ads12815 Aspergill
6	351	96.2	349	8	ADS12805	Ads12805 Aspergill
7	351	96.2	349	8	ADS12803	Ads12803 Aspergill
8	351	96.2	349	8	ADS12863	Ads12863 Aspergill
9	351	96.2	350	4	AAB82976	Aab82976 Aspergill
10	351	96.2	350	5	AAE15372	Aae15372 Aspergill
11	321	87.9	64	5	AAE15374	Aae15374 Aspergill
12	298	81.6	450	4	AAB82975	Aab82975 Trichoder
13	298	81.6	451	5	AAE15371	Aae15371 Trichoder
14	298	81.6	451	8	ADS12801	Ads12801 Trichoder
15	298	81.6	451	8	ADS12804	Ads12804 Trichoder
16	295	80.8	409	7	AB043144	Abo43144 A. thalia
17	295	80.8	409	7	ADB31925	Adb31925 Plant (A.
18	295	80.8	409	8	AD002271	Ado02271 Thalecres
19	292	80.0	64	5	AAE15373	Adouzzii inalecies Aael5373 Trichoder
20	172.5	47.3	84	5	ABP02534	Abp02534 Human ORF
21	172.5	47.3	200	8	ADS43437	Ads43437 Bacterial
22	172.5	47.3	230	2	AAW53806	Ads43437 Bacterial Aaw53806 Transcrip
23	172.5	47.3	230	8	ADT87049	Adt87049 Yeast Str
24	172.5	47.3	238	2	AAW53807	Aaw53807 Transcrip
25	171	46.8	68	5	AAE15382	Aae15382 Yeast HAC
26	171	46.8	68	8	ADS12859	Ads12859 Saccharom
27	123	33.7	168	5	AAU93013	Ads12039 Saccharom Aau93013 Arabidops
28	123	33.7	168	7	ADD30174	Add30174 Plant yie
29	123	33.7	168	8	ADI43893	Adi43893 Plant tra
30	123	33.7		3	AAG08861	Ad143893 Flant Cla
31	110.5	30.3	192	4	AAB82614	Aab82614 Maize roo
32	110.5	30.3	192	4	AAB82615	Aab82615 Maize roo
33	110.5	30.3	192	4	AAB82616	Aab82616 Maize roo
34	110.5	30.3	192	4	AAG66525	Aag66525 Maize roo
35	110.5	30.3	192	4	AAG66526	Aag66526 Maize roo
36	109.5	30.0	185	9	ADW17162	Adw17162 Eucalyptu
37	109	29.9	143	.3	AAB33151	Aab33151 Pinus rad
38	108	29.6	163	9	ADW17580	Adw17580 Pinus rad
39	105.5	28.9	170	8	ADM48147	Adm48147 Polypepti
40	101	27.7	261.	6	ABR47631	Abr47631 Breast ca
41	101	27.7	261	7	ADB75629	Adb75629 Prostate
42	101	27.7	261	8	ADL83138	Add73029 Prostate Adl83138 Human PRO
43	101	27.7	261	8	ADN16846	Adrio 3136 Human XB1
44	101	27.7	261	8	ADR14692	Adria646 Human NF-
45	101	27.7	261	8	ABM82329	Abm82329 Tumour-as
	201		201	9	111102323	ADMOZSZS TUMOUT-as

```
RESULT 1
ID
    AAB82977 standard; protein; 342 AA.
XX
AC
   AAB82977;
XX
DT
    11-SEP-2003 (revised)
```